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## (54) SULFONATED ELASTOMERS

We, Esso Research and Engineering Company, a Corporation duly organised and existing under the laws of the State of Delaware, United States of America, of Linden, New Jersey, United States of America, do hereby declare the invention for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the following state-

This invention relates to novel elastomers containing sulfonic acid groups or sulfonic acid salt moieties and to a process for preparing said elastomers. The term "elastomer" is employed in the specification and claims in its normal meaning of a polymer having rubber-like characteristics such as elasticity.

Polymers generally fall into two broad classes, thermoplastic and thermosetting resins. The thermoplastic resins may be readily worked by heating the polymer up to its softening point or melting point. They may then be processed by such deformation methods as vacuum forming, extrusion of a melt, compression molding, etc.

The thermoset resins can ordinarily not be reworked once they have hardened. In general, thermoset resins owe their unique properties to covalent crosslinks between polymer molecules. The crosslinks may be introduced, for example, by interaction of various monomers or other additives with or without heating.

Uncured elastomers such as natural rubber and butyl rubber are thermoplastic. They may, however, be crosslinked by vulcanization by the use of sulfur and accelerators which react with the carbon of the unsaturated bonds in the polymer molecules to form in effect a thermoset product which can no longer be fabricated or worked, except e.g. by machining. The vulcanized polymers have found wide utility because of the significant improvement in physical properties obtained by the crosslinking. Natural the significant improvement in physical properties obtained by the use of sulfur which reacts with the carbon of the unsaturated bonds in the polymer molecule to form a bridge between two molecules so that one polymer molecule is covalently bonded to the second molecule. If sufficient crosslinks of this type occur, all molecules are joined in a single giant molecule. Once crosslinked, the polymer is intractable and can no in a single genit angeotice. One curosumous, one popular is intractable and call no longer be fabricated except possibly by machining. It has, however, significantly improved physical properties. Thus, by vulcanizing rubber, elasticity, impact resistance, flexibility, thermostability and many other properties are either introduced or improved.

A third class of polymers, known as ionic polymers, has recently been developed which, although they are crosslinked, have a melt or softening point or softening range and may even be dissolved in various solvents. At normal use temperatures, these ionic polymers behave similar to crosslinked polymers. At elevated temperatures, however, they are readily deformed and worked in the same manner as thermoplastic resins. These ionic polymers (ionomers) owe their unique properties to the fact that crosslinking has been accomplished by ionic rather than covalent bonding between molecules of the polymer. Typical of these ionic polymers are copolymers of ethylene and ethylenically unsaturated mono- or dicarboxylic acids which have been neutralized by metal salts

(see, for example, British patent 1,011,981 and U.S. patent 3,264,272).



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2 1,268,264 2 Sulfonic acid ionmers have been prepared by copolymerization of a styrene sulfonic acid salt and other monomers to form plastic polymers containing ionic crosslinks, see, for example, U.S. patent 3,322,734. Methods of sulfonating polymers are well known to the art. For example, aromatic containing polymers are sulfonated by a method described in U.S. patent 3,072,618, wherein a complex of a lower alkyl phosphate and SO3 is used as the sulfonating agent. These sulfonated aromatic polymers have generally been sulfonated to a sufficient extent to be water-soluble in their acid form. Other aromatic containing resins have been sulfonated, converted to their alkali metal salts and used as ion exchange resins. 10 Water-soluble polymers have been prepared by reacting aromatic rings in styrenebutyl rubber graft polymers with SO, to form a viscous sulfonation product, see, for example, U.S.S.R. patent No. 211,079. 10 It has surprisingly been found that the physical properties of aromatic containing polymers may be improved by sulfunating the polymer to contain 0.2 to 21 mole % sulfonic acid groups. The tensile strength, Young modulus, and high temperature melt 15 strength may be further improved by converting at least a portion of the sulfonic acid 15 moietics to their corresponding amine or metal salts. According to the present invention there is provided a sulfonated elastomer containing monocyclic aromatic groups and 0.2 to 21 mole % sulfonic acid groups as 20 In particular the sulfonated elastomers of this invention can be prepared by (a) herein defined. 20 dissolving the unsulfonated form of the elastomer in a solvent which is also a solvent for the said complex of (b), and (b) reacting the elastomer at a temperature between -100° and 100° C with a sulfonating agent comprising (for example one to four moles of) sulfur trioxide and (for example one mole of) a complexing agent, selected from the Lewis bases containing oxygen, phosphorus or nitrogen, to introduce 0.2 to 21 mole % sulfonic acid groups into the elastomer. The sulfonated elastomer so formed can be 25 ionically crosslinked by neutralization with metal salts, amines or amine derivatives (as hereinafter specified whereby at least 1% of the sulfonic acid groups are in the 30 When employed in connection with the products and process of this invention, the term "aromatic" in "aromatic nuclei", "aromatic groups", "aromatic units", 30 means monocyclic aromatic. The expression "butyl rubber" as employed in the specification and claims means copolymers made from a polymerization reaction mixture having therein from 70 to 99.8% by weight of a C, to C, isoolefin, e.g. isobutylene, and 30 to 0.2% by weight 35 of a C, to C1, conjugated multiolefin, e.g. isoprene. The resulting copolymer contains 85 to 99.8% by weight of combined isoolefin and 0.2 to 15% of combined multiolefin. Buryl rubber generally has a Staudinger molecular weight of about 20,000 to about 500,000, preferably about 25,000 to about 400,000, especially about 100,000 to about 400,000; and a Wijs Iodine No. of about 0.5 to 50, preferably 1 to 15. The prepara-tion of butyl mober is described in U.S. patent 2,356,128. 40 For the purposes of this invention, it is preferred that the butyl rubber has incorporated therein from about 0.5 to 6% of combined multiolefin, more preferably 49 The first step in the preparation of sulfonated butyl rubber elastomers and corres-0.5 to 3%, e.g. 2%. 45 ponding ionomers of this invention is the grafting of styrene to the butyl rubber backbone. The grafting is accomplished by halogenation of the butyl rubber and subsequently reacting cationically polymerizable monomers (i.e. styrene) in the presence of an aluminum alkyl or haloaluminum alkyl catalyst. The graft polymer is then 50 reacted with a complex of sulfur trioxide with a Lewis base in order to sulfonate the 50 aromatic ring. The elastomer can then be converted to an ionomer by neutralizing the sulfonic acid moiety. Halogenated butyl rubber is commercially available and may be prepared by halo-55 genating butyl rubber in a solution containing between 1 to 60%, by weight of butyl rubber in a substantially inert C.-C. hydrocarbon solvent such as pentane, hexane, 55 heptane, etc., and contacting this butyl rubber cement with a halogen gas for a period of about 25 minutes whereby halogenated butyl rubber and a hydrogen halide are formed, the copolymer containing up to one halogen atom per double bond in the copolymer. The preparation of halogenated butvl rubbers is old in the art, see e.g. 60 U.S. Patent 3,099,644. This invention is not intended to be limited in any way by the 60 manner in which butyl rubber is halogenated and both chlorinated and brominated

butyl rubber are suitable for use in this invention.

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Illustrative of commercial halogenated butyl rubber is a chlorinated butyl rubber having about 1.25 wt. % chlorine, 2 mole % unsaturation and a viscosity average molecular weight of about 357,000.

The preparation of butyl rubber-styrene graft polymers from halogenated butyl rubber is disclosed, Belgian patent 701,850.

The structure of the ionomer former from a sulfonated butyl rubber-styrene graft polymer is thought to be:

where M+ is a metallic ion of Group I or, less preferably, of Groups II, III, IV, V, 10 VI-B, VII-B and VIII of the Periodic Table (hereinafter identified) of the

It will be readily evident that the technique disclosed is applicable to the formation of sulfonate ionomers from any elastomer containing an aromatic nucleus ring in its structure. A particularly suitable elastomer is homogeneous styrene isobutylene copolymer of the general structural unit:

The term "homogeneous" is used to distinguish this polymer from more conventional styrene-isobutylene copolymers which are in fact predominantly polymer blends of polystyrene and polyisobutylene.

Such homogeneous polymers may be prepared by solution polymerization under steady-state continuous polymerization conditions in a well-mixed tank type reactor. Solvent for the polymerization is either hexane or a hexane rich blend of hexane with methyl chloride. The polymer may contain as little as 0.5 mole % styrene or as much as 90 mole % styrene.

Other clastomers which may be sulfonated by the process of this invention include copolymers of styrene and butadiene, styrene and isoprene, chloroprene and styrene, copanyates of systeme and outstancing styrene and isoprene, currotypicto and styrene and acrylinities and styrene (SAN), styrene and acrylic acid, graft polymers of styrene onto polybutadiene, graft polymers of styrene-acrylonitrile copolymers (SAN) onto polybutadiene (ABS), styrene-isobutylene-isoprene terpolymers, etc.

A typical sulfonation procedure comprises contacting the polymer at about 0 to 100°C. for a period of a few seconds to several hours with a sulfonating complex which is preferably prepared by reacting about 2 to about 4 moles of sulfur trioxide with 1 mole of a phosphorous compound and recovering the sulfonated polymer.

In carrying out the present invention, an alkenyl aromatic-containing polymer is 35 sulfonated with a sulfonating agent comprising a sulfur trioxide donor in combination with a Lewis base complexing agent. In its preferred embodiment, the Lewis base 35

The term "sulfur trioxide donor" as used in the specification and claims means those substances including SO<sub>3</sub>, containing sulfur trioxide in a loosely bound form, which it can readily be liberated. Illustrative of the sulfur trioxide donors of this invention are sulfur trioxide, fuming sulfuric acid ("oleum", 20—30%), chlorosulfonic acid. The term "complexing agent" as used in the specification and claims means the Lewis bases suitable for use in the practice of

this invention.

The term "Lewis base" as used herein is used in its usual manner to define materials which donate electron pairs to form covalent bonds. It is intended to include mell-known Lewis bases as well as other complexing agents which act as Lewis bases

in the sulfonation reaction.

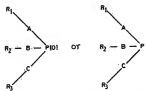
In a preferred embodiment, the base and sulfur trioxide donor are contacted to form a complex) prior to admixing them with the polymer. However, it is not essential to premix the sulfur trioxide donor with the complexing agent, but rather it is only necessary to have the latter compound present during the sulfonating reaction. For instance, the available sulfur trioxide donor and complexing agent may reaction. For instance, the available sulfur trioxide donor and complexing agent may in the sulfonating zone.

Because the complexing agent affects the reactivity of the sulfur trioxide

door, the sulfonation temperature need not be maintained below 0° C., but rather it may vary from as low as —100° C, to as high as 100° C. Moreover, the pressure is not a critical condition and may be adjusted to any satisfactory level. For instance, the sulfonation may be carried out from a reduced pressure of, say, 0.5 atmosphere up to a superatmospheric pressure in the area of 10 atmospheres. The most suitable conditions superatmospheric pressure in the area of 10 atmospheres. The most suitable conditions are temperatures of 15° to 40° C. and pressures which are approximately atmospheric. The sulfonation time will, of course, vary with the are approximately atmospheric. The sulfonation time will, of course, vary with the Generally, the reactions are complete within a few seconds to several hours after the Generally, the reactions are complete within a few seconds to several hours after the temperature and atmospheric pressure, the contact time should be 5 seconds to 25 or 30 minutes. Since the complexing agent reduces the activity of the sulfur trioxide donor, it is not necessary to limit the sulfonating time as is required in conventional

processes.

Illustrative of Lewis bases suitable for use as complexing agents are certain phosphorous compounds. While the phosphorous compound may be either inorganic or organic, it is preferred that the phosphorous compound be an organic compound having the general formula:



wherein A, B and C are each independently oxygen, or —CH<sub>2</sub>—; and R, Re R, are each independently selected from the group consisting of C, to C, alkyl, aryl, alkaryl or aralkyl. Various organic phosphires, phosphinates, phosphinates, phosphorates, phosphorates phosphorates and phosphines may be used as the complexing agent.

Illustrative of phosphorus containing inorganic complexing agents are phosphoric acid, phosphorus acid, pyrophosphoric acid, metaphosphoric acid, phosphonic acid and phosphinic acid. In addition to the acids, their moto, di- and tri-substituted derivatives may also be employed. However, the preferred phosphorous compounds are the trialkyl phosphates and phosphires.

Illustrative of other organic phospherous compounds suitable for use as complexing Illustrative of other organic phospherous compounds suitable for use as complexing agents are: riethyl phosphate, trimethyl phosphate, tripropyl phosphate, tri-butyl phosphate, tri-trithyl phosphite, etc.

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Other organic compounds suitable for use as complexing agents include the organic pyrophosphates. These compounds have the general formula:

#### R.R.R.R.P.O.

wherein R<sub>10</sub>, R<sub>20</sub>, R<sub>4</sub> and R<sub>7</sub> are each independently H, C<sub>7</sub> to C<sub>12</sub> alkyl, aryl, aralkyl or alkaryl, with the proviso that at least two of the groups are not H. The groups has used in the specification the term "substituted derivatives thereof" when such in reference to these R<sub>1</sub> to R<sub>2</sub> groups means nitro or halogen-substituted R groups.

Illustrative of such organic psychosphares are terrarelyll prophers.

Hlustrative of such organic pyrophosphates are tetrachyl pyrophosphate, tetramethyl pyrophosphate, dimethyl diethyl pyrophosphate and bis (2.4-dichlorophenyl) diethyl pyrophosphate. Where pyrophosphates are employed, as many as 13, bur more perferably 12 moles of sulfur trioxide donor may be cumplexed with each mole of blosphorous commonad

Other Lewis bases suitable as complexing agents are those Lewis bases which contain oxygen or nitrogen.

The nitrogen containing Lewis bases which form active complexes with sulfur trioxide donors and which are suitable for sulfonation of the unsaturated polymers of this invention have the general formulae:

wherein R<sub>1</sub>, R<sub>2</sub>, R<sub>3</sub>, and R<sub>32</sub> are each independently hydrogen, or C<sub>4</sub> to C<sub>34</sub> alkyl, aryl, alkaryl or aralkyl, with the proviso that where R<sub>4</sub> and R<sub>4</sub> are hydrogen, R<sub>16</sub> must not be hydrogen and R<sub>11</sub> is selected from the group consisting of C<sub>5</sub> to C<sub>34</sub> alkylene. Broadly speaking, the nitrogen containing Lewis base may be any primary, secondary or tertiary organic amine or a cyclic organic amine. Where the R<sub>35</sub> R<sub>15</sub>, R<sub>17</sub>, R<sub>11</sub> or R<sub>12</sub> radicals containing oxygen, chlorine, nitro groups or mixtures thereof. That is, R<sub>4</sub>, atoms, nitro groups or comprise both ethereal, or be substituted by halogen atoms, mixture of R<sub>15</sub>, R<sub>15</sub>, R<sub>16</sub>, R<sub>17</sub>, R<sub>18</sub>, R<sub>18</sub>, R<sub>18</sub>, R<sub>18</sub>, R<sub>19</sub>, R<sub>19</sub>, R<sub>19</sub>, R<sub>19</sub>, R<sub>10</sub> and R<sub>17</sub>; G these formulae, the term "substituted analogues" as used in the specification means groups as described and abstituted with

the substitutes referred to above. For the purposes of this invention, where R<sub>1</sub>, R<sub>2</sub>, R<sub>1</sub>, and R<sub>2</sub>, are ethereal, they shall be considered to fall in the class of "substituted analogues". In cases where R<sub>2</sub>, R<sub>3</sub>, and R<sub>3</sub>, contain aryl, alkaryl or aralkyl groups, the use of excess suffur trioxide donor is required to compensate for any sufformation of the aromatic units.

Illustrative of these nitrogen containing Lewis bases are trimethylamine, triethylamine, dimethylamline, diethylamline, piperidine, morpholine, N-ethyl morpholine, diethylaminoacetal, and tri 2-chloroethylamine.

Other suitable complexing agents containing nitrogen may be represented by the formula:

wherein  $R_{17}$ ,  $R_{18}$ ,  $R_{29}$ ,  $R_{14}$  and  $R_{12}$  are hydrogen, halogen, or  $C_4$  to  $C_{46}$  alkyl, aryl, alkaryl or aralkyl. In cases where  $R_{17}$ ,  $R_{19}$ ,  $R_{19}$ ,  $R_{14}$ , and  $R_{12}$  contain aromatic units, the use of excess sulfur trioxide donor is required to compensate for any sulfonation of these units. The organic  $R_{17}$ ,  $R_{12}$ ,  $R_{12}$ ,  $R_{14}$ , or  $R_{1}$  radicals may contain halogen or oxygen atoms, or nitrog groups. With reference to these  $R_{15}$  to  $R_{17}$  groups, the term "substituted analogues" as used in the specification means groups as described and substituted which oxygen or halogen or containing oxygen in the carbon chain. In addition, fused cyclic ring and polymeric structures are satisfactory.

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Illustrative of this type of compound are pyridine, 2-methyl pyridine, 2,6dimentally pyridine, quinoline, quinoline, and poly-2-vinyl pyridine.

The preferred nitrogen containing complexing agents are triethylamine and tri-n-

propylamine. The molar ratio of sulfur trioxide donor to nitrogen in the complex may be as high as 5 to 1, but the preferred ratio is 1 to 1.

Suitable solvents for the preparation of these complexes are chlorinated hydrocarbons. The preferred concentration of complex in solution is from 5 to 20% by weight, more preferably, 10 to 15%. The complexes may also be prepared by direct addition of the reagents if precautions are taken to dissipate the heat evolved. The reactions of complexes of SO<sub>3</sub> with molecules containing nitrogen and the polymer backbones containing aromatic nuclei have been found to be nonquantitative. Therefore,

the use of excess complex is desirable in such reactions. Oxygen containing Lewis bases have been found useful for complexing sulfur trioxide donor to facilitate reaction with the unsaturation of polymer molecules. The general formulae of such bases are:



wherein  $R_{11}$  and  $R_{19}$  are each independently  $C_2$ — $C_{16}$  alkyl, aryl, alkaryl or aralkyl and  $R_{01}$  is  $C_2$ — $C_{16}$  alkylene;  $R_{09}$  may be a substituted alkylene; the substituents may be  $G_1$  to  $G_2$  alkyl, aryl, alkaryl or aralkyl, For  $R_{13}$ ,  $R_{10}$  or  $R_{24}$  greater than  $G_{25}$  be  $G_1$  to  $G_2$  alkyl, aryl, alkaryl or aralkyl, For  $R_{13}$ ,  $R_{10}$  or  $R_{24}$  greater than  $G_{25}$  be  $G_1$  are  $G_2$  greater than  $G_2$  be  $G_3$  and  $G_4$  greater than  $G_2$  be  $G_3$  and  $G_4$  greater than  $G_2$  be  $G_3$  and  $G_4$  greater than  $G_2$  greater than  $G_3$  greate to R<sub>18</sub>, R<sub>19</sub> and R<sub>200</sub> the term "substituted analogues" as used in the specification means R11, R19 and R20 substituted as described above. Also suitable as oxygen containing Lewis bases are dioxane and substituted dioxanes. The presence of aryl, alkaryl or aralkyl groups in the base demands the use of excess SO, since some of the SO, is

used in sulfonation of the aromatic groups. Illustrative of the oxygen containing Lewis bases suitable for use in the practice of this invention are tetrahydrofuran, para-dioxane, 2,3-dichloro-1,4-dioxane, meta-dioxane, 2,4-dimethyl-1-1,3-dioxane, 2 phenyl-1,3-dioxane and bis(2-dichlorethyl) ether.

Other oxygen containing Lewis bases suitable as complexing agents are esters having the general formula:

wherein R21 and R22 are C1 to C14 alkyl, phenyl or benzyl or their nitro- or halosubstituents. Illustrative of such esters are benzyl acetate, butyl acetate, butyl propionate, methyl benzoate, hexyl acetate, isobutyl benzoate, ethyl-o-bromobenzoate, pnitrophenyl acetate, ethyl-n-butyrate, ethyl stearate and ethyl phenyl acetate.

The molar ratio of SO<sub>4</sub> donor to base may be as high as 15 to 1; preferably about 9:1 to about 1:1, more preferably about 5:1 to about 1:1, most preferably

about 2:1 to about 4:1, e.g. 3:1.

The preferred solvents for preparation of the complexes of sulfur trioxide donor with complexing agents containing oxygen are chlorinated hydrocarbons or the oxygen containing complexing agent. Illustrative of such chlorinated solvents are carbon tetrachloride, dichloroethane, chloroform and methylene chloride. The complexes may also be prepared by direct addition of reagents if precautions are taken to dissipate evolved

The reactions of ethereal complexes of SO<sub>3</sub> with the polymer chains containing heat. aromatic groups has been found to be nonquantitative. Therefore, the use of excess complex is desirable to give the required amount of sulfonation.

Any of the phosphorus, oxygen or nitrogen containing Lewis bases may be used individually as complexing agents or mixtures thereof may be used. The preferred

complexing agents are the organic phosphorous compounds. Where the phosphorus containing complexing agent is used, the complex may contain 1 to 15 moles of sulfur trioxide donor per mole of complexing agent, preferably 1 to 9 moles, more preferably 1 to 5 moles, most preferably 2 to 4 moles, e.g. 3 moles.

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The preferred complexing agents are trimethyl phosphite, triethyl phosphite and triethyl phosphate, more preferably triethyl phosphate. The preferred sulfur trioxide

In the practice of this invention, the polymer to be sulfonated is dissolved in a suitable solvent and reacted with a complex comprising the complexing agent and the sulfur trioxide donor at a temperature between -100° C and 100° C. The solvent medium must be a mutual one for the polymer and the complex. The solvent is

Illustrative of such solvents are alkanes, chlorinated alkanes, ethers, esters, or mixtures thereof. The alkane may be linear, branched or cyclic.

Illustrative of these alkanes are hexane, pentane, butane, cyclohexane, heptane and homologues thereof and analogues thereof. Illustrative of the chlorinated alkanes are methyl chloride, ethyl chloride, dichloroethane, chloroform, methylene chloride, carbon tetrachloride or any higher alkane or chlorinated alkane,

Illustrative of the suitable ethers and esters are tetrahydrofuran, p-dioxane, diethyl ether, amyl ethyl ether, bis-pentachloroethyl ether, bis- $\beta$ -chloroisopropyl ether, butyl acetate, isoamyl acetate and cyclohexylacetate,

Preferably, the solids content of the polymer solution is less than 25 wt. %. More preferably, 5 to 20 wt. %, most preferably 10 to 20 wt. %. Though cements having solid contents as high as 60 wt. % may be used there is a handling problem

The complex may be formed at -100° C. to +100° C., more preferably -40 to +40° C. Preferably, the complex is prepared immediately prior to use. In situ preparation of the complex in the polymer solution has also been found to be adequate.

Where the complex is prepared prior to use, its concentration in solvent should be 0.5 to 25 wt. %, more preferably 1 to 20 wt. %, most preferably 10 to 20 wt. %, e.g.

18 wt. ½.

Sulfonation of the polymer is conducted at a temperature between -100 and +100° C. Sulfonation occurs when the complex in solution is added to the polymer solution. The reaction time may be from 5 seconds to 3 hours. The product remains soluble throughout the reaction period. The product is best recovered by flashing off the solvents in hot water. The water also decomposes the unreacted complex.

The sulfonated polymers may be purified further by kneading in the presence of low boiling ketones or alcohols. The preferred materials for this purpose are acctone and methanol. After kneading, the polymers are dried on a hot mill. The recovered products are soluble in a variety of solvents, which indicates that the sulfonation has been accomplished without crosslinking,

Illustrative of the sulfur trioxide complex formed by reacting a complexing agent with SO, is the trialkyl phosphate/SO, complex shown in equation (1):

$$R = 0$$
  
 $R = 0$   
 $R = 0$ 

wherein R is an alkyl group and X represents the moles of SO<sub>5</sub> per mole of phosphorus compound. Preferably, X is about 1 to about 7; more preferably, about 2 to about 4, e.g. 3.

A typical sulfonation reaction proceeds as follows, R being an alkyl group:

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The metal ions which are suitable in forming the ionic copolymers of the present invention may be classified into two groups; non-complexed metal ions and complexed metal ions. In the non-complexed metal ion, the valence of the ion corresponds to the valence of the metal. These metal ions are obtained from commonly

known and used metal salts. The complex metal ions are those in which the metal is bonded to more than one type of salt group and at least one of which is ionized and one of which is not. Since the formation of ionic polymers requires only one ionized valence, such complex metal ions are equally well suited for use in the present invention. The utility of complex metal ions employed in the formation of ionic copolymers corresponds in their ionized valences to those of the non-complexed ions. The monovalent metals are, of course, excluded, but the higher valent metals may be included depending upon how many metal ions are complexed and how many can be ionized. The preferred complex metal ions are those in which all but one of the metal valences are complexed and one is readily ionized. In particular, the mixed salts of very weak acids such

as stearic acid and ionized acids such as formic and acetic acids may be utilized. The non-complexed metal ions which are suitable in forming ionic copolymers in the present invention comprise mono, di, tri and tetravalent metals in the Groups I, 

Fe+3, and Y+3. Suitable tetravalent metal ions are Sn+4, Zr+4, Ti+4, and Pb+4. In addition to the metal ions, other basic materials such as primary, secondary and tertiary amines may be to form the ionic bonds. The preferred amines have a basicity constant, Kin greater than 10-3. The basicity constant may be defined by the expres-

# $K_b = \frac{[Product]}{[\sim SO_3H] [Amine]}$

[Product] = Concentration of Product ~SO<sub>3</sub>H] = Concentration of Sulfonic Acid groups in the polymer [Amine] = Concentration of amine

The preferred amines have a K<sub>b</sub> value in the range of 10<sup>-3</sup> to about 10<sup>+6</sup>, preferably about 103 to 10-3. Illustrative of such amines are anhydrous piperazine, tri-npropylamine, triethylamine and triethanolamine. The secondary and tertiary amines are most preferred, especially piperazine and tri-n-propylamine.

The melt fabricability of the ionic polymer is affected not only by the number of crosslinks but to a much larger extent by the nature of the crosslink. For example, in the preparation of sulfonic acid ionomers of homogeneous styrene-isobutylene copolymers, the ionic content, where a monovalent salt [M+] is used is preferably 1 to 20 moles of [M+ styrene sulfonate] groups per 100 monomer units in the polymer; more

preferably, 4 to 16 moles; most preferably 8 to 12 mole. In a sulfonation process in accordance with this invention, the SO<sub>3</sub>-complex is present in the reaction medium at 0.2 mole to 21 moles per 100 repeating units in the polymer; preferably 1 to 20 moles; more preferably 2 to 16 moles; most preferably 4 to 12 moles. The sulfonated elastomers of this invention contain 0.2 mole % to 21 mole % -SO<sub>3</sub>H in the elastomer; more preferably 2 to 16 mole %; most preferably 4 to 12 mole %. From 1 to 100% (numerically) of the -SO<sub>2</sub>H groups may be

Throughout the specification and claims the term " mole % sulfonic acid groups " neutralized to form the ionomer.

means moles sulfonic acid groups per 100 monomer (i.e. repeating) units.

For divalent metal ions [M+2] such as Zn<sup>4</sup>2, Ca<sup>2</sup>2, Cu<sup>2</sup>2, and Ba<sup>2</sup>2, the ionic For a valent menu ions [m. ] such as Zu.; Ca.; Cu.; and Ma., the such as Zu.; Ca.; Cu.; and Ma.; the such as Zu.; Ca.; Cu.; and Ma.; the such as Zu.; Ca.; Cu.; and Co.; the monomer units. For trivalent metal ions [M. 13] such as Al 15, Fe 15, and Co 15, the monomer units. For trivalent metal ions [M. 13] such as Al 15, Fe 15, and Co 15, the

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	ionic content should preferably be less than 1.0 mole of [M <sup>+3</sup> (styrene sulfonate) <sub>5</sub> ] groups per 100 monomer units.	9
5	The distinct advantages of the sulfonated elastomers, and corresponding ionomers, of this invention may be more readily appreciated by reference to the following examples, of which Example 1 is a comparative example.	
	EXAMPLE 1	
10	A homogeneous styrene-isobutylene copolymer was prepared in a conventional overflow reactor. The reactor contained a central chamber housing a stirrer and two outer chambers for circulation of reactants. The reactor was suspended in a temperature bath. Two feed lines were used. The first contained the monomers dissolved in a solvent and the second contained dissolved catalyst. Both feeds were pre-chilled to the proper temperature.	ı
15	The reaction temperature was -90° C and the reaction time was 30 minutes. The monomer feed comprised 40 weight % styrene and 60 weight % isoburylene. The monomers were diluted to a 17.5 % concentration in a 60/40 (by vol.) mixture of methyl chloride and hexane. The catalyst was AlCl, at 0.12 wt. % concentration. After a reaction time of 30 minutes, the grant leaf of the catalyst was AlCl, at 0.12 wt. % concentration.	1
20	ing in isopropyl alcohol. The copolymer was recovered and dried. The intrinsic viscosity was 1.099 (1 g./liter of botten 25.84° C.). The copolymer had a viscosity average molecular weight of 330,000; a syrene content of 39.3 weight % was determined by nuclear magnetic resonance spectroscopy.  The product displayed poor stress bearing capacity, as shown in Table I, and was soluble in certain viscosity.	20
	but did not swell in polar solvents such as alcohol or water as evidenced in Table II.	
25	Paraman 2	
30	The product of Example 1 was sulfonated by the method disclosed in U.S. patent 3,072,618. Twenty grams of polymer was dissolved in 760 ml. of dry dichloroethane to which was added a 100 ml. solution of 3.35 ml. triethyl phosphate with 2.48 ml. of freshly distilled sulfur trioxide in dichloroethane. The reaction time was 10 minutes at 10° C. The polymer contained 7.12 × 10 <sup>-4</sup> equivalents of SO <sub>2</sub> H per gram (i.e. 4.9 moles of sulfonated styrene per 100 monomer units in the copolymer). The physical properties of the copolymers are shown in Table 1 and its solubility characteristics in Table 11.	30
	Example 3	
35	The product of Example 2 was converted into an ionic interpolymer by milling into it exact equivalents of sodium methoxide. On heating methyl alcohol is driven off. The material is easily processable, moldable, and re-workable. The strength properties are good (Table). The material displays solvency properties similar to those of the product described in Example 2 (see Table II).	35
40	Example 4  The material of Example 1 was sulfonated (using the general procedures outlined in Example 2) to obtain a product with 1.93 x 10 <sup>-3</sup> equivalents of SO <sub>2</sub> H/gm. of polymer, as determined by iteration (see Exemple 2).	40
45	polymer, as determined by titration (see Example 2) and elemental sulfur analyses. This corresponds to 13.3 moles of sulfonated styrene units per 100 monomer units. The material exhibits excellent stress supporting properties (see Table 1) and a broad temperature range over which it is rubbery. It is easily re-processable under molding conditions (280° F. and 12 tons pressure). The material has increased resistance to non-polar solvents and decreased resistance to polar solvents (Table II).	45
50	EXAMPLE 5  The material of Example 1 was sufformered by the method of Example 2 to obtain a product with 18.1 moles of sufformered supresse units per 100 monomer units. The product was very strong and elastic (Table 1) and re-processable under molding conditions. It was somewhat resistant to non-polar solvents but was greatly affected by polar solvents (see Table II).	50

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#### EXAMPLE 6

The product of Example 5 was converted into an ionic interpolymer by milling in exact equivalents of sodium methoxide. The product was difficultly processable. Apparently, an upper limit for ionic character was barely exceeded.

The material of Example 5 was converted into an ionic interpolymer by the addition of ZnO such that 5 moles of sulfonated styrene/100 monomer units were made ionic. The product was easily processable and re-processable at molding temperature (280° F.) and pressure (12 tons). The product was quite strong (Table I) and had good non-polar solvent resistance (Table II).

#### EXAMPLE 8

The material of Example 1 was sulfonated by the method of Example 2 to obtain a product with 20.7 moles of sulfonated styrene units per 100 monomer units. The product was strong (Table I) and re-processable. The material was swollen somewhat by non-polar solvents and was greatly affected by polar solvents (Table II).

#### EXAMPLE 9

The material of Example 8 was converted into an ionic interpolymer by adding ZnO such that 10 moles of sulfonated styrene/100 monomer units were made ionic. The product was easily processable and reprocessable under molding conditions. The material was only slightly swollen by non-polar solvents. (Table II).

Aftig 1.

Strength Properties of Sulfonated, Homogeneous Syrene-Isobutylene Copolymers at 25°C.

(Pull rate = 4°Vmin.)

	Material* (Example)	Tensile Strength lb./in.*	Modulus (300%) lb./in. <sup>2</sup>	Elongation %
-i	Styrene-Isobutylene Copolymer	275	50	1900
ci	4.9 mole % ~ SO <sub>3</sub> H	850	179	1100
'n	4.9 mole $\% \sim SO_3-Na^+$	750	390	750
₹	13.3 mole % ~ SO <sub>3</sub> H	1470	993	200
10	18.1 mole % ~ SO <sub>3</sub> H	1350	467	580
7.	13.1 mole $\% \sim SO_3H \div 5$ mole $\% Zn (\sim SO_3^-)_2$	1000	420	210
00	20.7 mole % ~ SO <sub>3</sub> H	930	463	440
o.	10.7 mole % ~ SO <sub>3</sub> H + 10 mole % Zn (~ SO <sub>3</sub> <sup>-</sup> ) <sub>2</sub>	1440	1100	380

\* Unfilled.

TABLE II

Behaviour of Sulfonated Homogeneous Styrene-Isobutylene Copolymers Toward Polar and Non-Polar Solvents (At 25°C.)

Volume Swell %

	Material (Example)	Benzene	Carbon Tetrachloride	Methanol	Watder
-	1. Styrene-Isobutylene Copolymer	solubie	soluble	0	0
7	4.9 mole % ~ SO <sub>3</sub> H	soluble	soluble	226	17.3
6	4.9 mole % ~SO <sub>3</sub> -Na+	soluble	923	4.2	3.3
4;	13.3 male % ~ SO <sub>3</sub> H	170	307	soluble	97.4
หก่	18.1 mole % ~ SO <sub>3</sub> H	13.9	50.8	soluble	191.6
7:	13.1 mole % ~SO <sub>3</sub> H <sup>+</sup> 5 mole % Zn (~SO <sub>3</sub> -) <sub>2</sub>	22.0	62.1	soluble	170
တ်	20.7 mole % ~ SO <sub>3</sub> H	17.6	48.6	soluble	150.5
9.	10.7 mole % ~ SO <sub>3</sub> H +10.0 mole % Za (~SO <sub>3</sub> ) <sub>2</sub>	29.6	6,98	soluble	58.9

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#### EXAMPLE 10

Using the method of Example 2, styrene-isobutylene copolymer was sulfonated at various levels. The modulus was determined on sulfonated samples both before and after preparation of ionomers by the method of Example 3.

Referring to the two figures of the accompanying drawings, Fig. I shows the effect of hydrogen bonding and ionic bonds on modulus. The figure shows that as the mole % of SO<sub>3</sub>H groups increases, there is a slight increase in modulus.

By contrast, conversion of the product to ionomers radically increases the modulus. For example, the Figure shows a sample of styrene-isobutylene copolymer containing about 16.15 mole % SO<sub>2</sub>H converted to an ionomer with varying degrees of ionic crosslinking. Surprisingly, there is a radical change in modulus with ionic

Fig. 2 shows the temperature dependence of tensile strength of unsulfonated styrene-isobutylene copolymer as compared with the same product containing 16.15 mole % SO<sub>3</sub>H and varying degrees of ionic content.

Curve A shows the tensile strength of the unsulfonated copolymer as a function of temperature. At 20° C. the copolymer has a tensile strength of less than 375 lb./in.2. At less than 60° C., the tensile strength is virtually zero (0).

By contrast, copolymer containing about 16.15 mole % SO,H (Curve B) has a tensile strength at 20° C. in excess of 600 pd and nearly 500 psi at 100° C. Conversion of only about 4 mole % SO,H to the calcium ionomer (Curve C)

results in a product having a tensile strength in excess of 900 psi at 20° C. and about 550 psi at 100° C.

The advantages of using a monovalent ion to form the ionomer are shown in Curve D. With about 12 mole %, of the SO<sub>2</sub>H groups converted to the sodium salt, 25 the copolymer has a tensile strength about 1500 psi; however, at 100° C, the tensile strength is less than 400 psi. Hence, despite a high room temperature tensile strength,

the copolymer is readily processed at moderate temperatures.

It is evident from the foregoing examples that useful styrene-isobutylene elastomers may contain 4.9 to 20.7 mole  $^{\prime}$ , SO,H groups. As little as I mole  $^{\prime}$ , (SO<sub>3</sub>),M wherein n represents the valence of the metal ion, M, results in an ionomer of improved 30 properties. More preferably, the ionic content, i.e. (SO<sub>2</sub>)<sub>n</sub>M groups, is 2 to 18 mole %; most preferably 4 to 12 mole %.

EXAMPLE 11

Previously examples described the sulfonation of a purified homogeneous styreneisobutylene copolymer. It is not necessary to purify the product before sulfonation, as described in this example. The material of Example 1 was prepared again, except that the catalyst activity was not killed by quenching the reaction in isopropyl alcohol. Rather, the reaction liquor (methyl chloride/hexane=60/40 by vol.) was maintained at -40° C. and the styrene-isobutylene copolymer was sulfonated directly with the triethyl phosphate and sulfur trioxide reagents. The polymer was recovered and analyzed. The material contained 5.57 moles of sulfonated styrene/100 monomer units. The properties of the material were found to be identical in all respects to a carefully purified preparation (in dichloroethane).

It is readily evident that the ionic polymer of homogeneous styrene-isobutylene copolymer is a superior thermoplastic material suitable for use in fabrication by extrusion, injection molding, or compression molding. Because of its unusual behaviour toward solvents, the product finds utility in the fabrication of rubberized paints, alcoholic thickeners, flocculents, dye receptors and adherents to metals.

#### EXAMPLE 12

A styrene-chlorobutyl graft polymer was prepared using the method of British Patent 1,174,323. Specifically, 100 grams of chlorobutyl rubber HT-1066 was dissolved in a solvent system of 900 ml, of heptane and 1000 ml, propane at -50° C. 206.3 gm. of freshly distilled styrene was added and then 2.87 gm. of AlEt<sub>9</sub>Cl catalyst. The reaction proceeded for 90 minutes. At the end of this time the product was recovered and dried; the yield was 249.0 gm. The product was extracted with acetone to remove styrene homopolymer. The rest of the material was solubilized in MEK (methyl etnyl ketone) to separate the gelations precipitate. The MEK soluble fraction was recovered and dried. 5 gm. of this material was dissolved in 190 ml. of dichloroethane. The sulfonation procedure of U.S. Patent 3,072,618 was used then

0.38 ml. of triethyl phosphate was added followed by 0.28 ml. of sulfur trioxide at 5° C. The reaction proceeded for 10 minutes and was terminated by quenching, and the product recovered.

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Analysis of the styrene-butyl graft copolymer showed an intrinsic viscosity of 0.77 in toluene and 42% styrene by infrared spectroscopy. The properties of a compression molded pad were: tensile strength=2800 lb./in."; elongation at break=

Acid base titration of the ionic styrene-butyl graft copolymer in a 50/50 mixture of toluene and methanol with 0.1 N-standard KOC/CH<sub>0</sub>), showed that this product had 8.83 × 10 1 acid equivalents per gram. The sulfur content was found to be 1840 0.05 × 10 data equaterins per grain. The suntir content was found to be 2.84%, by weight. The product thus contained 6.5 miles // sulfionic acid groups. The sulfonated styrene-buyl graft copolymer was made into the ionic product

by dissolving the material in a 50/50 mixture of toluene and methanol and adding 8.83 x 10- equivalents of sodium methoxide. The precipitated product was dried and compression molded at 300° F. and 12 tons of pressure for 10 minutes. The molded object was thermoplastic and could be remolded under the previous conditions. The properties of a molded pad were tensile strength=1370 lb./in.2 and clongation at break -225%. The ionic graft copolymer was decidedly stiffer than the non-ionic graft

The ionic graft copolymer displayed unusual solvent resistance compared to the copolymer. non-ionic graft copolymer. A summary of the effects of various solvents is shown in Table III. It is seen that while the non-ionic graft copolymer was highly swollen or dissolved in certain organics, the ionic graft copolymer was insoluble or was merely

# swollen.

### TABLE III Behaviour of Graft Copolymer and Ionic Graft Copolymer Toward Solvents

Solvent	Graft Copolymer	Ionic Graft Copolymer
Benzene	soluble	swells slightly
Carbon tetrachloride	soluble	swells
Heptane	highly swollen	insoluble (not swollen)
Methanol	insoluble	insoluble
Acetone	insoluble	insoluble
Methyl Ethyl Ketone	soluble	insoluble
Water	insoluble	insoluble

#### EXAMPLE 13

A commercial styrene butadiene rubber containing 23 wt. % styrene and 77 wt. % butadiene having a membrane osmometry number average molecular weight of 120,000 was sulfonated in the following manner. The SBR was purified by dissolving 50 grams of rubber in 500 ml. of hexane

and precipitating the polymer in 10 liters of methanol. The rubber was dried on a

rubber mill at about 240° F. Five gram quantities of the purified material were sulfonated using an (EtO), PO: (SO<sub>a</sub>)<sub>2</sub> complex to form polymers having various levels of sulfonation.

Sulfonation was carried out in solvent mixtures of cyclohexane and methylene chloride. The polymer was dissolved in the solvent mixture and the complexes which were prepared in methylene chloride was added dropwise with stirring. The reaction was carried out over a ten (10) minute period at 25° C.

Conversion of the sulfonic acid containing polymers to ionomers was accomplished by reacting the sulfonic acid polymer with twice the stoichiometric quantity of NaOH dissolved in methanol. The polymers were recovered by dropping the polymer solution into boiling water, thereby flashing off the solvents and precipitating the polymer.

The recovered polymer was dried at 130° C.

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Table V summarizes the various experimental results. It is noted that the sulfonic acid polymer is soluble at 6.5 mole percent SO<sub>3</sub>H which indicates there has been no crosslinking. The polymers were thermoplastic and could be molded and remolded at 130° C. and 9000 psig. Although sulfonation has been described in terms of sulfonation of the aromatic 5 nucleus, it is thought that in polymers such as SBR which contain a high degree of 5 non-aromatic unsaturation, the sites of unsaturation are also sulfonated. Not wishing to be bound by the theory, it is postulated that sulfonic acid moieties are distributed between sites of non-aromatic unsaturation and aromatic rings dependent on the 10 mole ratio of the unsaturation to aromatic ring and the reactivity of the sites sulfonated. The industrian of the unsaturation to anomatic ring and the reactive than the sizes of unsaturation are more reactive than the aromatic ring. Hence, in a polymer such as SBR the preponderance of the sulfonic acid groups are 10 added at non-aromatic unsaturation sites. Since the SO<sub>3</sub>H groups may be pendant either from the aromatic ring or the 15 polymer backbone, their concentration in the polymer is expressed in mole % based polymer to account their concentration in the polymer is expressed in more polymer after than the aromatic units. It is desirable that the sulfonance SRR of this invention contains 0.2 to about 20 mole % SO<sub>2</sub>H groups; preferably about 12 mole % mole %; more preferably about 0.5 to about 12 mole % abo 15 suitably 0.48 to 6.5 mole %, most preferably about 1 to about 8 mole %, e.g., 3.0 mole %. From about 1 to 100 % of the SOoH groups may be neutralized to form the 20

TABLE V

Preparation and Properties of a Sulfonated Styrene-Butadiene Copolymer (SBR)

Condition of Reaction	soluble						
Mole % SO <sub>3</sub> Na	0	0.50	0	1.51	0	2.90	0
Mole % SO <sub>3</sub> H	0.48	0	1.42	0	2.82	0	6.5
ті. Сн <sub>3</sub> он	0	1.0	0	1.0	0	5.0	0
gm. NaOH	0	0.0328	0	0.0984	0	0.1968	0
CHO	1.0	1.0	1.0	1.0	2.0	2.0	3.0
Moles (BtO) <sub>3</sub> PO Used	1.37 × 10-4	1.37 > 10-4	$4.1\times10^{-4}$	4.1 × 10-4	$8.2 \times 10^{-6}$	$8.2\times10^{-4}$	1.92 × 10-3
Moles SO <sub>3</sub> Used	4.1 × 10-4	4.1 × 10-4	$1.23 \times 10^{-3}$	$1.23\times10^{-3}$	2,46 × 10-3	$2.46 \times 10^{-3}$	$5.76 \times 10^{-3}$
Moles Reactivity Available in Solution	0.0823	0.0823	0.0823	0.0823	0.0823	0.0823	0.0823
ml. Solvents	100 ml. cyclohexane + 40 ml. CH <sub>2</sub> Cl <sub>2</sub>	100 ml. cyclohexane + 40 ml. CH <sub>4</sub> Cl <sub>2</sub>	100 ml. cyclohexane ⊹ 40 ml. CH <sub>2</sub> Cl <sub>2</sub>	100 ml. cyclohexane + 40 ml. CH <sub>2</sub> Cl <sub>2</sub>	100 ml. cyclohexane + 40 ml. CH <sub>2</sub> Cl <sub>2</sub>	100 ml, cyclohexane + 40 ml, CH <sub>2</sub> Cl <sub>2</sub>	100 ml. cyclohexane + 40 ml. CH <sub>2</sub> Cl <sub>2</sub>
SBR.	2	6	NO.	10	ĸ	w	3

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The improvement in physical properties is readily demonstrated by comparing styrene-butadiene copolymer with the same polymer both sulfonated and converted to the sodium ionomer. Physical properties are shown in Table VI.

TABLE VI Properties of Sulphonated SBR

Mole % SO <sub>3</sub> H	Mole % SO <sub>3</sub> Na	Tensile lb/in²	Modulus (50%) lb/in²	Modulus (30%) lb/in²	% Elong.	% Set at Break
0 0.48	0	35	39	41	570	400
0.48	ő	75	40	63	680	350
0 1.42	0.5 0	64	40	53	800	350
1.42	ŏ	150	60	95	700	200
0	1.51	230	90	170	630	40
2.82	0	460	81	230	580	30
0	2.90	530	156	375	650	5
6.5	0	650	170	600	340	0

It is readily evident that sulfonation improves properties of the polymer. For example, with as little as 0.48 mole % SO<sub>2</sub>H in the polymer, the tensile strength has been increased more than 100%. The sulfonated polymers may be cured in any

#### EXAMPLE 14

Using the method of U.S. Patent 32.65,765, a styrene-bundiene block polymer spreared containing 30 wt. % styrene (S) and 70 wr. % bundiene (B). The structure of the polymer was S—B—S wherein S represent styrene sequences having a number average molecular weight of about 15,000 and styrene-sens bundiene sequences having number average molecular weight of about 15,000 and Sprearest bundiene sequences having number average molecular weight of about 15,000 and sprearest bundiene sequences having number average molecular weights of about 51,000 and 51,000 and

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TABLE VII

Sulfonation of a Styrene-Butadiene-Styrene Block Copolymer

gm. Polymer	Moles Reactivity Available in Solution	Solvent System	Moles SO <sub>a</sub> Used	Moles (EtO) <sub>3</sub> PO Used	MI. CICH2CH2CI	Mole % SO <sub>3</sub> H
10	0.158	100 ml. cyclohexane + 40 ml. CH <sub>2</sub> Cl <sub>2</sub>	1.59 × 10 <sup>-3</sup>	1.59 × 10-3	2.0	0.45
10	0.158	100 ml. cyclohexane + 40 ml. CH <sub>2</sub> G <sub>2</sub>	$6.34 \times 10^{-3}$	$6.34 \times 10^{-3}$	5.0	0.81
10	0.158	100 ml, cyclohexane + 40 ml, CH <sub>2</sub> Cl <sub>2</sub>	$1.27 \times 10^{-3}$	$1.27 \times 10^{-2}$	10.0	1.24

Example 1 were The syrane-isobunylene copolyments propared in the manner of Example 1 were sulforated using complexing agents other than the phosphorous containing Lewis beas. The lev IIII shows the results of sulforation using p-dioxane, triethylamine 5 has detrainfunduran as the complexing agent. It is readily evident that these Lewis bases are effective complexing agents.

TABLE VIII
Sulfonation of Styrene-Isobutylene Copolymers

			5	Complex Preparation			
Polymer Styrene Content (mole %)	mg	ml. Solv. (Carbon Terrachloride)	Moles SO <sub>3</sub>	Moles Complexing Agent	Ml. Solvent	- % Sulfur in Product	Mole % SO <sub>3</sub> H
(Mn = 175,000)	10	200	0.0144	0.0144 p-dioxane	10 CH <sub>2</sub> Cl <sub>2</sub>	2.84	5.28
(Mn = 175,000)	10	200	0.0144	0.0144 (Et) <sub>3</sub> N	15 CH2CI2	0.29	0.54
$(\overline{Mm} = 110,000)$	10	200	0.0313	0.0313 THF	20 CH2Cl2	0.26	99.0
$\frac{50.0}{(Mn = 110,000)}$	10	200	0.0313	0.0313 (Et) <sub>3</sub> N	20 CH2CI2	0.15	0.38

	All of the products formed were soluble in the reaction medium. The products	
	All of the products formed were soluble in the reaction methods formation of were found to have improved physical properties even without formation of were found to have improved polymer containing 1.24 mole % SO <sub>3</sub> H was	
	were found to have improved physical properties even without the ionomer. For example, sulfonated polymer containing 1.24 mole % SO <sub>3</sub> H was the ionomer. For example, sulfonated physical properties were attributed	
	the ionomer. For example, sulfonated polymer contaming the contamination of the contamination	5
5	more resistant to flow in a cold press. The improved physical projects to hydrogen bonding. The product may be readily converted to the ionomer by the	
,	method of Example 14.	
	WHAT WE CLAIM IS:-	
	1 A culforated elastomer containing monocyclic aromatic groups and	
	mole % sulfonic acid groups as herein defined.	10
10	mole % sulfonic acid groups as herein defined.  2. A sulfonated elastomer as claimed in claim 1, containing 0.2 to 16 mole %	
10	sulfonic acid groups.	
	sulfonic acid groups.  3. A sulfonated elastomer as claimed in claim 2, containing 0.2 to 12 mole %	
	sulfonic acid groups.	
		15
15	4. A vulcanizable suironated classrolle contained in any preceding claim.  having a sulfonic acid group content as claimed in any preceding claim.	
10	5. A vulcanizable suifonated classomer conding of claims 1 to 3.	
	5. A vulcanizable suifonated classonier comparising of claims 1 to 3. having a sulfonic acid group content as claimed in any of claims 1 to 3. having a sulfonic acid group content as claimed in claim 5, containing 0.48	
	6. A vulcanizable sunbilated clastomer as distance	
	to 6.5 mole % sulfonic acid groups.	20
20	7. A vulcanizable suitonated elastomer complained in any of claims 1 to 3.	
	polymer having a sulfonic acid group content as claime a great polymer of a styrene-	
	polymer having a sulfonic acid group content as claimed in any objects.  8. A vulcanizable sulfonated elastomer comprising a graft polymer of a styrene-	
	acrylonitrile copolymer on polybutaniene, the grant polybutaniene	
	group content as claimed in any of claims 1 to 3.	25
25	9. A vulcanizable sulfonated elastomer comprising a gamup content as claimed in	
	<ol> <li>A vulcanizable sulfonated elastomer comprising a grant population on polybutadiene, the graft polymer having a sulfonic acid group content as claimed in</li> </ol>	
	any of claims 1 to 3.	
	10. A sulfonated homogeneous stylene-isotraty end of claims 1 to 3.	
	10. A sulfonated homogeneous synthesistoty of claims 1 to 3, having a sulfonic acid group content as claimed in any of claims 1 to 3, 11. A sulfonated elastomer as claimed in any preceding claim and substantially	30
30	11. A suitonated elasionici as channed in any p	
	as herein described.  12. A sulfonated elastomer as claimed in any of claims 1 to 11 and substantially	
	12. A sulfonated elastomer as claimed in any of claims 1, 5, 8 and 12 to 15. as herein described with reference to any one of Examples 2, 4, 5, 8 and 12 to 15.	
	as herein described with reference to any one of Examples 2, 3, 3, 3 and any of claims 1  13. A process for preparing a sulfonated elastomer claimed in any of claims 1	35
	13. A process for preparing a statement	22
35	to 10, which comprises:  (a) dissolving the unsulfonated form of the elastomer in a solvent, which is also	
	a solvent for the said complex of (b), and	
	a solvent for the said complex of (b), and (b) reacting the elastomer at a temperature between -100° C and 100° C (b) reacting the elastomer at a temperature by reacting a Lewis base containing	
	(b) reacting the elastomer at a temperature between with sufficient of an SO <sub>3</sub> complex, prepared by reacting a Lewis base containing with sufficient of an SO <sub>3</sub> complex, prepared by reacting a Lewis base containing with sufficient of an SO <sub>3</sub> complex, prepared by reacting a Lewis base containing	40
		40
40	required sulfonic acid group content into the clastomer,	
	required sulfonic acid group content into the classonic.  14. An ionomer comprising a sulfonated elastonic as claimed in any of claims the interest of the comprising a sulfonic acid groups have been neutralized to	
	14. An ionomer comprising a sulfonated classifier a sulfonate of to 10 in which at least 1% of the sulfonic acid groups have been neutralized to 1 to 10 in which at least 1% of the sulfonic acid groups have been neutralized to	
	1 to 10 in which at least 1% of the suitonic acid groups have a primary, secondary form the corresponding amine or metal salt, wherein the amine is a primary, secondary form the corresponding amine or metal salt, wherein the amine is a primary, secondary form the corresponding amine or metal salt, where the primary is at least one selected from Groups I to V, VIB,	45
	form the corresponding amine or metal sait, wherein the amine and the view of tertiary amine and the metal is at least one selected from Groups I to V, VIB, or tertiary amine and the metal is at least one selected from Groups I to V, VIB,	13
45	or tertiary amine and the metal is at least one strength of the Elements herein identified.  VIIB and VIII of the Periodic Table of the Elements herein identified.	
	VIIB and VIII of the Periodic Table of the Elements Herein Idealm 18, 2011.  15. An ionomer as claimed in claim 14, wherein the metal is sodium, potassium, 15. An ionomer as claimed in claim 14, wherein the metal is sodium, potassium, 15. An ionomer as claimed in claim 14, wherein the metal is sodium, potassium, 15. An ionomer as claimed in claim 14, wherein the metal is sodium, potassium, 15. An ionomer as claimed in claim 14, wherein the metal is sodium, potassium, 15. An ionomer as claimed in claim 14, wherein the metal is sodium, potassium, 15. An ionomer as claimed in claim 14, wherein the metal is sodium, potassium, 15. An ionomer as claimed in claim 14, wherein the metal is sodium, potassium, 15. An ionomer as claimed in claim 14, wherein the metal is sodium, potassium, 15. An ionomer as claimed in claim 14, wherein the metal is sodium, potassium, 15. An ionomer as claimed in claim 14, wherein the metal is sodium, potassium, 15. An ionomer as claimed in claim 14, wherein the metal is sodium, potassium, 15. An ionomer as claimed in claim 14, wherein the metal is sodium, potassium, 15. An ionomer as claimed in claim 14, wherein the metal is sodium, potassium, 15. An ionomer as claimed in claim 14, wherein the metal is sodium, potassium, 15. An ionomer as claimed in claim 14, wherein the metal is sodium, 15. An ionomer as claimed in claim 14, wherein the claim 15. An ionomer as claimed in claim 14, wherein the claim 15. An ionomer as claimed in claim 15. An	
	15. An ionomer as claimed in claim 14, which is thereof. calcium, copper, zinc, nickel, barium, aluminium or mixtures thereof.	
	calcium, copper, zinc, nickel, barium, aluminum of Indian substantially as described	

16. An ionomer as claimed in claim 14 or claim 15 and substantially as described

17. An ionomer as claimed in claim 14 or claim 15 and substantially as described

herein with reference to any one of Examples 3, 6, 7, 9, 12 and 13.

K, J, VERYARD,

15 Suffolk Street,

herein.

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FIG.1.
INFLUENCE OF HYDROGEN BONDS AND IONIC BONDS ON THE MODULUS

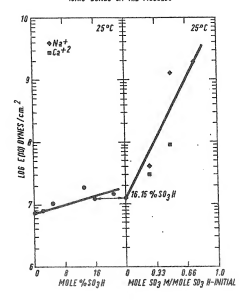


FIG. 2.

TEMPERATURE DEPENDENCE OF

TENSILE STRENGTH FOR

HYDROGEN BONDED AND IONIC INTERPOLYMERS

